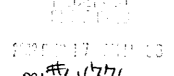
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#### **BIODEGRADATION STUDY REPORT**

## Revision 1

Wednesday, November 06, 2002

# Biodegradation Screen Study for Telomer-Type Alcohols

## **PROJECT NUMBER**

3M Projects ID: **E01-0684**Pace Contract Analytical Projects ID: **CA085** 

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# **PROJECT DATES**

Project Initiation: August 9, 2000 Project Completion: September 18, 2001

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# **Executive Summary**

A screening study was undertaken to determine whether the fluorochemical telomer intermediate (telomer alcohol) biodegrades when exposed to municipal wastewater treatment sludge. The study included the preparation of cultures for a six sample-point comparative study, and included test cultures, blank cultures, and control cultures. The microbial inoculum for cultures was sludge obtained from the Twin Cities Metro Wastewater Treatment Facility. Telomer alcohol test substance was added to cultures at approximately 5 μM concentrations. Cultures were incubated with shaking at 25°C. Solid phase extraction of cultures for recovery of biodegradation products and parent analytes was employed. An HPLC/MS analytical method was developed for analysis of the telomer alcohols and expected products. The HPLC/MS analysis of culture extracts provided strong evidence that the telomer alcohols were biodegraded. The observed loss of telomer alcohols occurred concomitant with the appearance of several expected perfluorinated carboxylic acids. Unexpectedly, transiently formed polyfluorinated  $\beta$ -oxidation intermediates were observed. The  $\beta$ -oxidation pathway was suspected to be the major route of biodegradation resulting in the observed even-numbered carbon chain length perfluorinated fatty acids. Minor end products, as odd-numbered carbon chain length perfluorinated fatty acids, were also observed and were likely the products of fatty acid α-oxidation, a minor pathway of fatty acid metabolism usually observed in branched-chain fatty acids. After the 16day test period, perflourinated carboxylic acids ranging from C5 to C12 were observed in the test cultures. These compounds were not in controls or blank cultures. Perfluorooctanoate (PFOA), was the only compound quantitatively analyzed and, based on mass balance data accounted for approximately 6-7% of the total telomer alcohols initially present in the test cultures.

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# 1.0 Project Personnel

1.1	Spons	sor Company	3M
1.2	Spons	sor Representative	Dr. James K. Lundberg
1.3	Contra	act Facility Personnel:	
	1.3.1	Study Director	Dr. Cleston C. Lange
	1.3.2	Laboratory Management:	Mr. Bruce E. Warden
	1.3.3	Sample Preparation	Ms. Angela L. Schuler
			Dr. Cleston C. Lange
	1.3.4	HPLC/MS Analyst	Ms. Angela L. Schuler
	1.3.5	Sample Custodian:	Dr. Cleston C. Lange
	1.3.6	Report Author	Dr. Cleston C. Lange

# 2.0 Data Requirements and Revision Justification

The sponsor representative desired an aerobic biodegradability screening study to be conducted using the telomer alcohol mixture as test substrate. The study was initiated on August 9, 2000 as a non-GLP study.

The report issued Friday, November 16, 2001 is revised to substitute "Dupont Zonyl BA type Telomer, or Zonyl type Telomer" with Telomer. The reason for the revision is to more accurately reflect the impact of results to the general class of telomer alcohols.

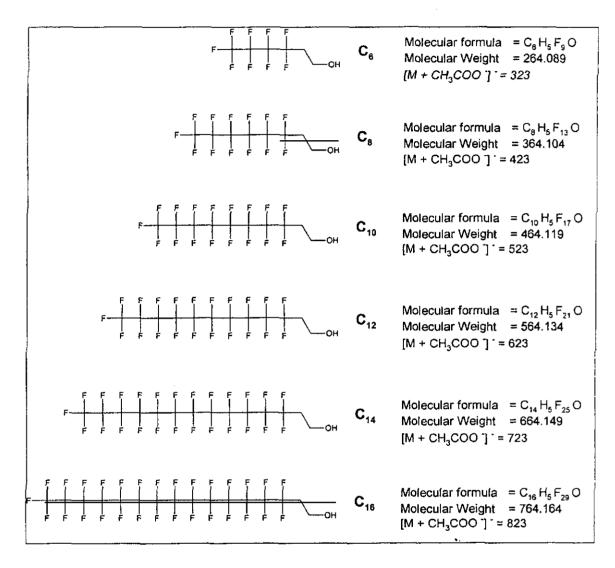
## 3.0 Project Objective

This study was conducted in order to elucidate whether the fluorotelomer intermediate (telomer alcohol) is biodegradable under aerobic conditions using a microbial rich inoculum of municipal wastewater treatment sludge. This study was similar to earlier studies conducted for 3M by Pace as Pace projects CA058 <sup>1</sup> and CA097 <sup>2</sup> and CA104 <sup>3</sup>, CA105 <sup>4</sup>, and CA132 <sup>5</sup>. The development of an analytical method and the analysis of parent analytes and possible products were critical parameters for the determination of biodegradation.

## 4.0 Test Article

The test article used for this study was the Zonyl BA-type telomer alcohol fluorochemical surfactant intermediate. Two sources of test material were used for this study. The first source was provided by the sponsor company, 3M, as approximately 2.5 g of crystalline solid test material labeled Zonyl BA-N telomer alcohol, 3M tracability number TN-A-2186, on July 7, 2000. The sponsor did not provide an MSDS, a chain of custody, the purity information for the test article, nor an expiration date for the material. This material was given a test, control, and reference (TCR) number at Pace as CA-TCR02-009 and was stored at 4°C. Material CA-TCR02-009 was used primarily for method development purposes of the study, including the first set of cultures prepared and original LC/MS method development.

A commercial vendor, Aldrich Chemical Company, provided the second source of telomer alcohol used. Material was purchased as Zonyl BA-L fluorotelomer intermediate (1999 Aldrich Catalog number 42,151-0), also called perfluoroalkylethanol. The material characteristics were boiling point 145-245°C, F(CF<sub>2</sub>)<sub>n</sub>CH<sub>2</sub>CH<sub>2</sub>OH where n equals approximately 7 to 8, and molecular weight was reported as M<sub>n</sub> ca. 443.70 wt. An expiration date and the purity, or percent composition, of the material were not provided. The purchased material was received at Pace on September 22, 2000. Upon receipt at Pace, the material was given tracability number CA-TNC00-254 and was stored at 4°C, or less. A representative chemical compilation of the Zonyl BA-type telomer alcohol is shown in Figure 1.



**Figure 1.** The telomer alcohols. Based on the HPLC/MS average peak area response observed in the six abiotic control cultures, the composition of the test material was  $6.4\% \pm 0.3\%$  as  $C_6$  telomer alcohol,  $39.3\% \pm 1.5\%$  as  $C_8$ ,  $27.7\% \pm 2.1\%$  as  $C_{10}$ ,  $17.5\% \pm 1.2\%$  as  $C_{12}$ ,  $7.2\% \pm 1.0\%$  as  $C_{14}$ , and  $2.0\% \pm 0.2\%$  as  $C_{16}$ .

# 5.0 Reference Materials

The sponsor provided reference material. The neat material was stored at -80°C.

NS: Expiration date not specified

# 5.1. Perfluorooctanoate Ammonium Salt (PFOA)

Purity: 95.2 %; 3M#: TCR-99131-37; Pace #: CA-TCR02-001; Expires: NS



Figure 2. Perfluorooctanoate ammonium salt

# 6.0 Receipt/Generation of Samples

Samples were not received, but were generated as an inherent part of this study. All of the experimental cultures prepared for each culture set were extracted by solid phase extraction (SPE) methodology to generate three analytical samples per culture. The analytical samples, as SPE eluates collected for each culture, were labeled as SPE eluates 1, 2 or 3. Of the three eluates generated for each sample, eluate 1 was the 25 mL aqueous sample eluate and eluates 2 and 3 were 25 mL methanol eluates from the SPE cartridge. Qualitative and semi-quantitative HPLC/MS analysis was conducted for eluate 2 only, and the resulting data was evaluated to determine whether biodegradation occurred.

## 7.0 Methods

# 7.1 Sample Preparation

# 7.1.1, Collection of Sludge.

The sludge for this study was obtained from the Twin Cities Municipal waste treatment facility in a manner consistent with the sludge collected for other studies listed in section 3.0. The sludge used for the preparation of the first set of cultures was collected on July 31, 2000 and

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1.55

delivered with Pace chain of custody (COC) form 465254. The sludge for the second set of cultures was collected on September 18, 2000 and was accompanied with COC # 528791. Sludge used for the second set of cultures in this study was also used in preparation of cultures for other fluorochemical biodegradation studies (CA105 <sup>4</sup> and CA132 <sup>5</sup>), and was shown to be active for biodegradation of other fluorochemical compounds in those studies.

To prepare cultures, sludge was obtained from the primary municipal waste treatment facility in the Twin Cities area. Arrangements were made for Pace personnel to retrieve fresh mixed liquor suspended solids (MLSS) from the aeration units at the Twin Cites Metro Wastewater Treatment Facility located in St. Paul, MN. Typically, Four liters of MLSS was collected by Pace laboratory personnel and delivered as four 1-liter Nalgene polypropylene bottles containing MLSS, and were accompanied with a corresponding chain of custody with date collected. Upon receipt at Pace Science Solutions, the individual bottles were labeled #1 through #4. The suspended solids in the bottles were allowed to settle at least 24 hours at 4°C ± 3°C. The settled solids "sludge" were then used to prepare MLSS plus sludge for use in preparing test cultures. A sludge characterization analysis was not conducted as part of this screening study.

The settled sludge in each bottle constituted approximately 20% of the volume, or approximately 200 mL volume in a 1-liter bottle, based on visual observations-and was consistent with observation of earlier sludge collections.

7.1.2. Culture Preparation.

The first set of cultures were prepared August 9, 2000 and incubation of

those cultures continued with intermittent culture harvests until the final

harvest on September 13, 2000, for a total incubation time of 35 days.

The second set of cultures were prepared on September 27, 2000 and

incubation of those cultures continued with intermittent culture harvests

until the final harvest on October 12, 2000, for a total of 16 days

incubation. The culture preparation procedure described below was

used, and was documented as Pace standard operating procedure

(SOP) CAG-SP-03 6.

Cultures were prepared using a mineral salts medium defined by EPA

Guideline OPPTS 835.3200. The mineral salts medium pH was 7.4 and

contained per liter, 0.334 g Na<sub>2</sub>HPO<sub>4</sub>-2H<sub>2</sub>O, 0.005 g NH<sub>4</sub>Cl, 0.2175 g

K<sub>2</sub>HPO<sub>4</sub>, and 0.085 KH<sub>2</sub>PO<sub>4</sub>, 0.0275 g CaCl<sub>2</sub>-anhydrous, 0.0225 g

 $MgSO_4$ -7 $H_2O$ , and 0.00025 g FeCl<sub>3</sub>-6 $H_2O$ .

Two liters of a mineral salts medium containing 100 mL of settled sludge

was prepared and contained 1 mL of methanol. To each test culture was

added 25 mL of this mineral medium containing sludge.

A mineral salts medium, un-sterilized and without sludge, was prepared.

This mineral salts medium without sludge was used to prepare the 25 mL

no-sludge (abiotic) control cultures.

All cultures were prepared by dispensing 25 mL of appropriate mineral

salts medium solution into clear sterile 125 mL Nalgene polycarbonate

culture flasks containing labels with appropriate identification information.

Note: The mineral salts medium that contained sludge had to be swirled

regularly during dispensing in order to keep the mixture homogenous

and prevent the sludge from settling out of the solution.

The test substance, either Zonyl BA-N telomer alcohol for culture set one

(Stock ID CA058-SS-004 at 10,280 µg/mL in methanol) or Zonyl BA-L

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telomer alcohol for culture set two (Stock ID CA085-SW-001 at 7,288  $\mu$ g/mL in methanol), was added to the test cultures by transferring 5  $\mu$ L and 8 $\mu$ L, respectively, to the appropriate test cultures. The final test concentration in cultures was 2.056  $\mu$ g/mL of Zonyl BA-N telomer alcohol for set one, and 2.332  $\mu$ g/mL of Zonyl BA-L telomer alcohol for set two. Blank control cultures received 25 mL of mineral medium solution, as did the test culture, but without addition of the test substrate.

All of the day zero cultures were prepared and immediately frozen at  $-20\,^{\circ}$ C for storage until SPE preparation could be conducted. All other cultures were placed in temperature controlled shaking incubators that were maintained at  $25\,^{\circ}$ C  $\pm$   $3\,^{\circ}$ C. Cultures were removed from the incubators at designated time points. Upon removal from the incubator, cultures were either immediately frozen, or immediately prepared for analysis by solid phase extraction.

All culture preparation information, including times, analyte additions, etc. were recorded in sample preparation worksheets and signed and dated by the preparation analyst. All original data sheets were maintained in project specific binder labeled as Project CA085.

# 7.1.3. Solid Phase Extraction of Cultures

The solid phase extraction procedure described below was documented as Pace standard operating procedure (SOP) CAG-SP-04 <sup>7</sup>.

All cultures and control cultures were prepared by solid-phase extraction methodology using SEP-VAC C18 6cc SPE cartridges from Waters Corporation (Part No. WAT036905). A sample label was applied to each SPE cartridge prior to use, and each cartridge was packed with plug of quartz glass wool to deter plugging. Each SPE cartridge was washed prior to use by drawing 5 mL of methanol and then 5 mL of aqueous 1% acetic acid solution through the cartridge. These wash solution eluates were discarded to waste. All of the SPE eluates for this study were collected in clear I-Chem vials with labels that identified them as eluate 1, 2 or 3, as defined below.

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Frozen cultures were thawed at ambient room temperature before extraction. Following thawing, and prior to solid phase extraction, 0.25 mL of glacial acetic acid was added to each of the cultures yielding a final concentration of 1% acetic acid. The content of each acidified culture was swirled to mix, and then drawn by vacuum through the appropriately labeled SPE cartridge by carefully pouring the contents of the culture flask into the SPE cartridge. The aqueous eluate was collected in an I-Chem vial labeled eluate 1, removed from the vacuum manifold, and capped. Then, 25 mL of methanol was added to the culture flask, the flask sealed, and vigorously shaken. The cap was then removed from the flask, and the methanol content (25 mL) drawn through the SPE cartridge, collected in an I-Chem vial labeled eluate 2. Eluate 2 was expected to contain a majority of the analyte that was in the original culture.

As a precaution that some analyte may be retained in the SPE cartridge or in the culture flask, a second 25 mL methanol eluate was collected in a similar fashion to that collected for eluate 2, and was labeled eluate 3. Aliquots of eluates 2 and 3 were transferred to autovials, capped, and then quantitatively analyzed by HPLC/MS. The remaining volume of each eluate was stored at 4°C ± 2°C.

The final SPE extractions of the day 7, 14 and 16 day cultures from set two occurred on September 24, 2001.

## 7.2 Instrumental Analysis (LC/MS)

# 7.2.1 Instrument Parameters

The HPLC/MS method used was a modified version of Pace method CAG-ORG-23 <sup>8</sup>, as described below.

Analysis of culture extracts from culture set one, prepared with Zonyl BA-N telomer alcohol, was conducted using instrument "STING" which included an HP1100 HPLC pump with Gilson 215 liquid handling system in line with a Micromass Quattro II triple-quadrapole mass spectrometer detector.

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Analysis of culture extracts from culture set two, prepared with Zonyl BA-L telomer alcohol, were analyzed both on instrument "STING" and instrument "10LCMS03" which consisted of a Waters 2690 HPLC system in line with a Micromass Quattro II triple-quadrapole mass spectrometer detector. Typical conditions for analysis of telomer alcohols and fluorochemical acid products were as follows:

Mass Spectrometer	<u>HPLC</u>		
lonization mode: API-ES negative			
Desolvation Temperature: 200°C	Time (min).	<u>%A</u>	<u>%B</u>
Source Block Temperature: 150°C	0.00	97.0	3.0
RF lens: 0.2	0.50	97.0	3.0
Extractor: 3	5.00	5.0	95.0
Cone: 8	11.00	5.0	95.0
Capillary: 3.50	11.50	97.0	3.0
LM Resolution: 14	15.00	97.0	3.0
HM Resolution: 14	Flow: 1 mL/min, s	plitter app	rox. 3:1
on Energy: 2.5	Solvent A= 2 mM	ammoniu	m acetate
_ens 6: 2	Solvent B= Metha	nol	
MS1 Multiplier: 650	Column Tempera	ture: ambi	ent
MS/MS: Collision Energy-varied from 10	to 40 V for individual	Experimer	nts,

Typical injection volumes for samples and calibration standards were 50  $\mu$ L. The 4.6 x 150 mm Betasil C8 column used for the quantitation of extracts from culture set two had serial number 1101567H and the 4 x 35 mm NG1 column used had serial number 15104. A pressure-regulated splitter was used with an approximate split ration of 2:1 (Waste:MS)

# 7.2.2 Qualitative Parent Analyte and Products Analysis.

Parent Analyte	Expected fons (m/z)
C <sub>8</sub> telomer alcohol acetate adduct	423
C <sub>10</sub> telomer alcohol acetate adduct	523
C <sub>12</sub> telomer alcohol acetate adduct	623
C <sub>14</sub> telomer alcohol acetate adduct	723
C <sub>16</sub> telomer alcohol acetate adduct	823

Table 1. Parent telomer alcohols analyzed.

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To conduct the analysis of parent compounds and expected biodegradation products, the HPLC/MS system was set up with the same chromatographic configuration as that described in method CAG-ORG-23 <sup>8</sup>. The method utilized two columns in tandem with flow to a pressure relief valve that serves as a flow through splitter to the mass spectrometer Z-spray source.

Specific MS conditions for mass analysis of the telomer alcohols and acid products were developed during this study. The mass spectrometer, Micromass triple-quadrapole mass spectrometer with Z-spray ion source, was operated with electrospray ionization in either selected ion-recording (SIR) mode or with mass-range (150-1000 m/z) scanning on MS1. Discrete chromatographic peaks with singly charged negative ions, (M + Acetate)<sup>-</sup> for parent telomer alcohols and (M - H<sup>+</sup>)<sup>-</sup> for expected acid biodegradation products, were observed and monitored:

Expected Product	Expected anion	Expected anions (m/z)
	structure	
Perfluorobutyrate (C <sub>4</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>2</sub> COO	213,169
Perfluoropentanoate (C <sub>5</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>3</sub> COO	263, 219
Perfluorohexanoate (C <sub>6</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>4</sub> COO	313, 269
Perfluoroheptanoate (C <sub>7</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>5</sub> COO	363, 319
Perfluorooctanoate (C <sub>8</sub> , PFOA)	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>6</sub> COO	413, 369
Perfluorononanoate (C <sub>9</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>7</sub> COO	463, 419
Perfluorodecanoate (C <sub>10</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>8</sub> COO	513, 469
Perfluoroundecanoate (C <sub>11</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>9</sub> COO	563, 519
Perfluorododecanoate (C <sub>12</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>10</sub> COO	613, 569
2H, 2H-perfluorooctanoate (C <sub>8</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>5</sub> CH <sub>2</sub> COO	377
2H, 2H-perfluorodecanoate (C <sub>10</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>7</sub> CH <sub>2</sub> COO	477
2H, 2H-perfluorododecanoate (C <sub>12</sub> )	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>9</sub> CH <sub>2</sub> COO	577

**Table 2.** Expected products, and expected anions for each.

# 7.2.3 Quantitative Analysis.

Quantitative analysis was conducted only for perfluorooctanoate (PFOA) and only on extracts of culture set two. Quantitative analysis was performed by the external standard method using SPE extracted calibration standards (eluate 2 only) and quadratic calibration curves. The HPLC/MS analysis of culture set one extracts was used primarily for method development purposes.

Typical injection volumes for samples and calibration standards were 50  $\mu$ L. The 4.6 x 150 mm Betasil C8 column used for the quantitation of extracts from culture set two had serial number 1101567H and the 4 x 35 mm NG1 column used had serial number 15104. The pressure-regulated splitter had no identifying number to distinguish it. The part number for ordering the pressure relief valve (splitter) was Alltech catalog part number 39025.

## 7.3 Data Transformations and Calculations

#### 7.3.1 Molar Calculations:

Because all data was collected on an ng/mL basis (part per billion, ppb), a transformation from ng/mL to molar concentrations had to be conducted to obtain mass balance information when applicable. The mole conversion values for each analyte are as follows:

PFOA molecular weight, as ammonium salt, is 431.10 Zonyl BA-L telomer alcohol average molecular weight is 434.70.

# 7.3.2 Conversion of ng/mL to micromolar (μM) and nanomolar (nM).(Working Examples):

2,232 ng/mL telomer alcohol =  $(2,232 \text{ ng/mL})*(1 \text{nmole} / 434.7 \text{ ng}) = 5.135 \text{ nmole/mL} = 5.135 \text{ µmole/liter} = 5.135 \text{ µM}—assuming 100% purity.}$ 

150 ng/mL PFOA (NH4+ salt) = (150 ng/mL) \* (1 nmole / 431.1 ng)

= 0.3479 nmole/mL = 0.3479  $\mu$ mole/L = 0.3479  $\mu$ M = 347.9 nM Page 16 of 38

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## 7.3.3 Molar Mass Balance Calculations (Theoretical Yield):

Convert all ng/mL values to their corresponding molar concentrations as  $\mu M$  or nM (see section 7.3.2, above). Divide the sum of the analyte concentrations by the known concentration of starting compound and represent the final result as a percentage of the known starting concentration.

# (Working Example):

If, the starting concentration of BA-L telomer alcohol was at  $5.135~\mu M$  And, after incubation, the following was determined:

if, PFOA was detected at 0.3479μM

Then, the mass balance is as follows:

Mass balance =  $[(0.3479 \mu M) / 5.135 \mu M] \times 100\%$ 

Mass balance =  $[0.0678] \times 100\% = 6.78\%$ 

Or, 6.78% of the telomer BA-L alcohol was oxidized to form PFOA.

#### 7.4 Software Versions

Microsoft™ Excel 2000 was used for data processing and producing tables. Microsoft™ Word 2000 was used for processing the analytical report text. Adobe Acrobat 4.0 was used for generation of the final electronic report. Masslynx version 3.2 was used for data collection and peak integration. ACD Chemsketch version 4.0 was used for chemical drawings.

8.0 Results

8.1 Sludge Characterization

A chemical analysis of the mixed liquor suspended solids used for the samples

prepared on August 9, 2000 was conducted, and characterization information

can be found in the final report for project CA058<sup>1</sup>. The sludge for samples

prepared on September 27, 2000 was not characterized, but was obtained from

the Twin Cities Municipal waste treatment facility in a manner consistent with the

sludge collected and used for project CA058.

8.2 Quality Control/Sample Matrix Spike Results.

The determination of the analyte recoveries from sample matrix spikes was not

included as part of this screening study. Recoveries are only reflected in the

ability to achieve molar mass balance based on expected parent and product

yields from samples and by the use of a sludge-extracted curve for semi-

quantitative analysis of PFOA.

8.3 Analytical Blanks

Methanol solvent blanks were injected onto the HPLC/MS column and

quantitatively analyzed to determine the instrument background analyte

concentration and carry over during the analysis.

The no-sludge (abiotic) controls containing mineral salts medium with test

analyte, and culture blanks, containing mineral salts medium with test analyte

and no sludge, were prepared and analyzed. Abiotic controls and blank cultures

were prepared and incubated in an identical manner to biodegradation test

cultures. The results were used to determine whether the sample matrix

contained any of the analytes of interest and whether biodegradation could be

attributed to the bioactivity of the sludge. Fluorochemical analytes were not

detected in blanks and biodegradation did not occur in abiotic controls.

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## 8.4 HPLC/MS Analysis Results

Semi-quantitative analysis was conducted only for the SPE eluate 2 extracts from the second set of cultures, prepared September 27, 2000, and were performed on September of 2001 as sequence CA085\_092401b.spl (with full scan MS data), and CA085\_092701a.spl (SIR data) on instrument 10LCMS03 in the Bio-Analytical services group at Pace. The qualitative HPLC/MS analyses and HPLC/MS method development for this study were conducted in September and October of 2000 as analytical sequence runs S091500.spl, S091800.spl, S092000.spl, S092100.spl, S092600.spl, S1000200.spl, and S100400.spl on Pace instrument "STING".

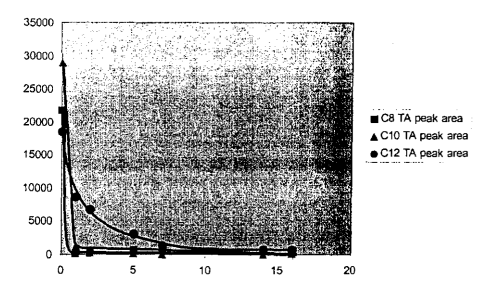
## 8.4.2 Qualitative HPLC/MS and HPLC/MS/MS Results

The first set of test cultures containing sludge and BA-N telomer alcohol substrate provided evidence that biodegradation occurred. The HPLC/MS data showed near total loss of all of the telomer alcohol peaks in the HPLC/MS chromatogram, with formation of chromatographic peaks that had mass spectra consistent with perfluorinated acids. However, at that time, the analytical test method was not complete. Concerns about the integrity of the test material resulted in a second set of test cultures being prepared using commercially purchased Zonyl BA-L telomer intermediate.

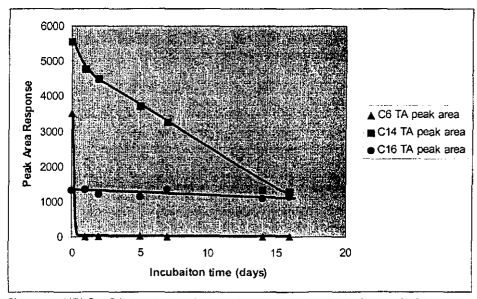
Although each sample, control, and blank for the second set of cultures was prepared and incubated in duplicate, analysis was conducted for SPE eluate 2 extracts from one sample, control and blank per time point.

The data showed that the sample test cultures underwent rapid loss of the  $C_6$ ,  $C_8$ ,  $C_{10}$ , and  $C_{12}$ -telomer alcohols, and moderate loss of the  $C_{14}$ -telomer alcohol over the 16 days of incubation, as determined by decreasing peak area response for each telomer alcohol ion at the sequential sampling time points (**Figures 3 and 4**). The  $C_{16}$ -telomer alcohol was degraded very slowly and showed little loss.

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**Figure 3.** HPLC/MS integrated peak areas for the most abundant telomer alcohols (TA). Near complete degradation of all three was observed. Data plotted with the best manual fit of the data.



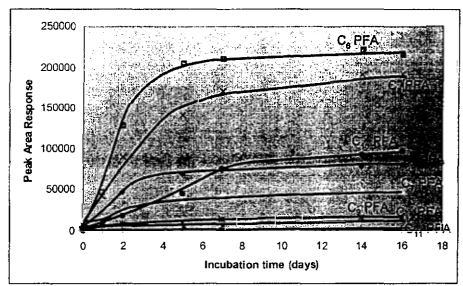
**Figure 4.** HPLC/MS integrated peak areas for the less abundant telomer alcohols (TA). Near complete degradation was observed for the  $C_6$ -TA. Moderate degradation of the  $C_{14}$ -TA was observed and little degradation of the  $C_{16}$ -TA was observed. Data plotted with the best manual fit of the data.

Concomitant with the loss in telomer alcohols was the formation of compounds with HPLC/MS ions and retention times consistent with perfluorinated acids (PFAs), including perfluoroctanoate (PFOA, C<sub>8</sub>-PFA), which was confirmed as an end product. The transformation of telomer alcohols to PFAs was rapid, with a large increase in all of the PFA peak area responses at day 1, and continuing increase through day-16 (Figure 5). No degradation to form PFAs was observed in the abiotic no-sludge controls.

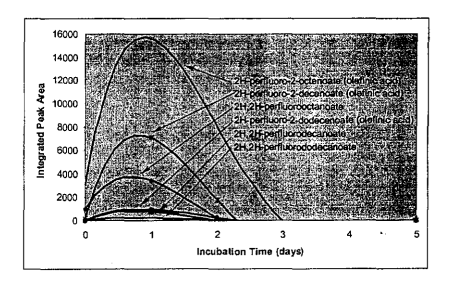
Although PFOA was accurately quantified, many other perfluorinated acids were qualitatively observed and their relative concentrations determined based on the observed increasing peak area response. The observed perfluorinated fatty acid end products were: perfluoropentanoic acid ( $C_5$  PFA), perfluorohexanoic acid ( $C_6$ -PFA), perfluorooctanoic acid ( $C_6$ -PFA), perfluorononanoic acid ( $C_9$ -PFA), perfluorodecanoic acid ( $C_{10}$ -PFA), perfluorodecanoic acid ( $C_{11}$ -PFA) and perfluorododecanoic acid ( $C_{12}$ -PFA).

Transiently formed intermediate compounds were also observed at early time points following the initial exposure of the sludge to the telomer alcohol substrate (**Figure 6**). The transient compounds were suspected to be: 2H, 2H-perfluorooctanoate; 2H, 2H –perfluorodecanoate; 2H, 2H Perfluorododecanoate and the possible  $\beta$ -oxidation pathway intermediates: 2H-perfluoro-2-octenoate, 2H-perfluoro-2-decenoate, and 2H-perfluoro-2-dodecenoate. **Table 3** shows the MS ions and retention times observed for the polyfluorinated and perfluorinated acid products and the telomer alcohol parent substrates.

HPLC/MS/MS Data (**Table 4**) support the identification of the suspected metabolites..



**Figure 5.** Integrated peak area responses for the perfluorinated fatty acids (PFAs) observed in test culture extracts. Even numbered carbon chain length carboxylic acids were the most abundant carboxylic acid peaks and were: perfluorohexanoate ( $C_6$ -PFA), perfluorooctanoate ( $C_8$ -PFA) and perfluorodecanoate ( $C_{10}$ -PFA).



**Figure 6.** Transient polyfluorinated fatty acid intermediates observed. The 2H-perfluorinated olefinic fatty acids are suspected β-oxidation intermediates that formed rapidly upon initial exposure to telomer alcohols, and then were transformed to their corresponding perfluorinated carboxylic acids end products

Analytes Observed	Chemical Formula	Retention Time	LC/MS anions	
			observed	
C <sub>6</sub> -telomer alcohol*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> OHCH <sub>3</sub> COO	7.63	323	
Ce-telomer alcohol*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>5</sub> CH <sub>2</sub> CH <sub>2</sub> OHCH <sub>3</sub> COO	8.13	423	
C <sub>10</sub> -telomer alcohol*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>7</sub> CH <sub>2</sub> CH <sub>2</sub> OHCH <sub>3</sub> COO	8.50	523	
C <sub>12</sub> -telomer alcohol*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>9</sub> CH <sub>2</sub> CH <sub>2</sub> OHCH <sub>3</sub> COO	8.96	623	
C <sub>14</sub> -telomer alcohol*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>11</sub> CH <sub>2</sub> CH <sub>2</sub> OHCH <sub>3</sub> COO	9.53	723	
C <sub>16</sub> -telomer alcohol*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>13</sub> CH <sub>2</sub> CH <sub>2</sub> OHCH <sub>3</sub> COO	10.25	823	
2H, 2H-perfluorooctanoate*	CF3(CF2)5CH2COO	7,17	377	
2H, 2H-perfluorodecanoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>7</sub> CH <sub>2</sub> COO	7.38	477	
2H, 2H Perfluorododecanoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>9</sub> CH <sub>2</sub> COO	7.49	577	
2H-perfluoro-2-octenoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>4</sub> CF=CHCOO	7.14	357,293	
2H-perfluoro-2-decenoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>6</sub> CF=CHCOO	7.35	457,393	
2H-Perfluoro-2-dodecenoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>8</sub> CF=CHCOO	7.52	557,493	
Perfluoropentanoate	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>3</sub> COO	6.62	263, 219	
Perfluorohexanoate	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>4</sub> COO	6.88	313, 269	
Perfluoroheptanoate	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>5</sub> COO	7.07	363, 319	
Perfluorooctanoate	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>6</sub> COO	7.22	413, 369	
Perfluorononanoate	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>7</sub> COO	7.30	463, 419	
Perfluorodecanoate	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>8</sub> COO	7.41	513, 469	
Perfluoroundecanoate	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>9</sub> COO	7.48	563, 519	
Perfluorododecanoate	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>10</sub> COO	7.52	613, 569	

<sup>\*;</sup> Only anions corresponding to even chain length molecules were observed.

Table 3. Parent and products anions observed in by HPLC/MS analysis of SPE eluate 2 extracts from test cultures containing sludge and telomer alcohol substrate. The parent anion signals decreased in test cultures over time concomitant with increases in product signals. Many of the telomer alcohol signals decreased below detection limits in test cultures.

Product Analytes Observed	Chemical Formula	Retention	LC/MS/MS	LC/MS/MS daughter
		Time	Parent anion	anions observed
2H,2H-perfluorooctanoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>5</sub> CH <sub>2</sub> COO	7.17	377	333,313,293
2H,2H-perfluorodecanoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>7</sub> CH <sub>2</sub> COO	7.38	477	413,393
2H,2H Perfluorododecanoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>9</sub> CH <sub>2</sub> COO	7.49	577	
2H-perfluoro-2-octenoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>4</sub> CF=CHCOO	7.14	357	313,293,243,143,119
2H-perfluoro-2-decenoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>6</sub> CF=CHCOO	7.35	457	413,393,343
2H-Perfluoro-2-dodecenoate*	CF <sub>3</sub> (CF <sub>2</sub> ) <sub>8</sub> CF=CHCOO	7.52	557	493

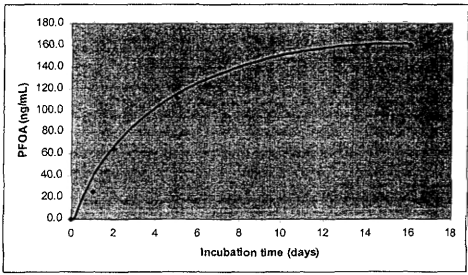
Table 4. HPLC/MS/MS data for suspected -oxidation products. Parent and daughter anions from the HPLC/MS/MS analysis of the SPE eluate 2 extract from the 1-day biodegradation sample (CA085-0927-SA-033 E2). The parent anions were observed as transiently formed anions in early sample point biodegradation samples only, with the most intense signals for the ions observed at day one. The MS/MS data collected for each product was consistent with the predicted chemical formula. Common mass loss from the parent anion, attributed to loss of CO<sub>2</sub> (mass 44) and HF (mass 20), were observed.

# 8.4.2 Quantitative Analysis of PFOA

The HPLC/MS multi-component calibration standards used in this study for product quantitation contained the target analytes: PFOA, PFOSulfinate, PFOS, FOSA, N-MeFOSA, M556, M570, and N-MeFOSE alcohol. Five calibration standards were prepared by addition of a known amount of perfluorinated analytes to a culture medium with sludge, and then extraction by solid phase extraction in manner identical to the treatment of samples. The PFOA was the only target analyte of that mixture that was used for this study. The exact concentrations of the standards were used in calibration curves for quantitative analysis.

All calibration standards and sample eluates were stored at 4°C in refrigerator ID 0213 until analysis. Aliquots were transferred to autovials and capped for use in HPLC/MS runs. The Instrument calibrations were performed for PFOA by use of a five point calibration with quadratic fit of the data. The LLOQ was 11.0 ng/mL PFOA, and the coefficient of determination (r) for PFOA was 0.965. Quantitative sequence runs contained calibration standards at the beginning and end of the run.

The biodegradation of the telomer alcohols resulted in an increase in the measured levels of PFOA during the 16-day study (Figure 7).



**Figure 7.** The measured perfluorooctanoate (PFOA) in cultures containing sludge and fluorotelomer alcohol. The PFOA was not measurable in cultures that did not contain sludge. Data plotted with the best manual fit of the data and concentrations shown were not adjusted for reference material purity of 95.2%.

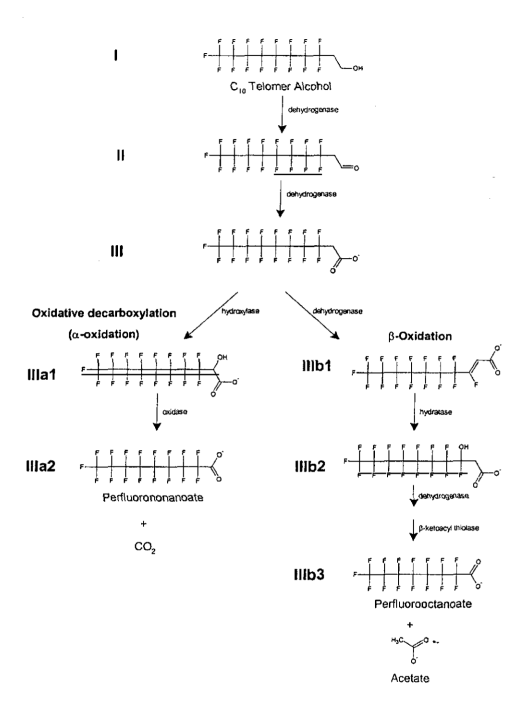


Figure 8. Proposed biodegradation pathway for telomer alcohols, shown for  $C_{10}$ -telomer alcohol as example. The β-oxidation pathway branch (IIIb) involves primary formation of a fatty acyl-CoA thioester that is not shown, to clearly depict the products observed by HPLC/MS. The fatty acyl-CoA thiol bond is readily hydrolyzed to yield the corresponding fatty acids observed.

## 9.0 Conclusions

This study has demonstrated that fluorochemical telomer alcohols are biodegradable and are transformed to perfluorinated acid end products. Although the telomer alcohols are supplied only as even numbered carbon chain compounds, the end product perfluorinated acids consist of both even number and odd number carbon chains, with even numbered carbon chain length acids predominating as the major end products. This observation suggests that, following the initial oxidation of the hydroxyl-carbon to form the primary carboxylic acid (telomer carboxylate), two oxidation mechanisms exist as shown in **Figure 8**, **III.** The first, less utilized route (**Figure 8**, **IIIa**), involves oxidation of the  $\alpha$ -carbon concomitant with decarboxylation (oxidative decarboxylation) to form an odd chain length perfluorinated carboxylic acid, which does not undergo further biotransformation. The second observed route of oxidation (**Figure 8**, **IIIb**) is the more common route of fatty acid  $\beta$ -oxidation, as evidenced by the formation of detectable transient  $\beta$ -oxidation polyfluorinated fatty acid intermediates and the more abundant even number carbon chain length carboxylic acids

Although toxicity of the telomer alcohols was not addressed as part of this study, the microbial degradation observed here over the entirety of the 16-day study suggests that microorganisms present in the sludge were not significantly inhibited by the tested concentration of the test material.

#### 8.0 Literature Cited

- 1.Final Reports for Pace Project CA058 (3M # E00-2252), "2-Week N-EtFOSE Alcohol Biodegradation Screen Study Report" and "Aerobic Biodegradation of N-EtFOSE alcohol Study Report". Author: Cleston C. Lange, Ph.D.
- 2. Final report for Pace Project CA097 (3M # E01-0415), "The 18-Day Aerobic Biodegradation Study of Perfluorooctanesulfonyl-based Chemistries." Author: Cleston C. Lange, Ph.D.
- 3. Final report for Pace Project CA104 (3M # E01-0444), "The 35-Day Aerobic Biodegradation Study of Perfluorooctanesulfonate (PFOS)." Author: Cleston C. Lange, Ph.D.
- 4. Final Report for Pace Project CA105 (3M # E01-0683), "Fluorochemical Adipate (L15468) Biodegradation Screen Study." Final report issued July 10, 2001. Study Director: Cleston C. Lange, Ph.D.
- 5. Final Report for Pace Project CA132 (3M # E01-0682), "The Aerobic Biodegradation Study of the Fluorochemical FC807-Diester". Final report issued May 29, 2001. Study Director: Cleston C. Lange, Ph.D.
- 6. Pace method CAG-SP-03. "Culture Preparation for Assessment of Aerobic Biodegradability of Fluorochemicals Using Municipal or Industrial Sludge as Microbial Inoculum."
- 7.Pace method CAG-SP-04. "C18 Solid Phase Extraction Procedure for Fluorochemicals Recovery from Aqueous/Sludge Matrices."
- 8. Pace method CAG-ORG-23 "Quantitative Analysis of Fluorochemicals by High Performance Liquid Chromatography with Mass Spectrometric Detection."

9.0 Sample and Data Retention

At a minimum, one copy of all pertinent raw data and one copy of the signed final report will be retained in the Pace Analytical-Tier 2 data archives for a minimum period of 2 years after completion of the project. The remaining sample extracts will be retained at the Pace Analytical facility for a period of 2 years after completion of the project at 4°C in

the Carroll walk-in cooler (Pace ID 0140) located in the Pace Analytical-Tier 2 facility.

The following will be provided to the sponsor (3M):

The original signed analytical report and one copy of the signed original.

The final scanned report (read only) and pertinent electronic data on a CD.

 All original Data, correspondence, chromatograms, sample & standards prep sheets, etc.

Upon request before 2 years, the stored samples may be sent to the sponsor.

All electronic copies of the instrumental raw data will be archived onto CD disks and one copy provided to 3M and one copy retained at Pace.

Facility data will be retained for a period of 10 years. Facility data is available for inspection and includes the following records:

Training records

Controlled storage temperature logs

· Standard preparation logs

Calibration and maintenance logs

Chemical and solvent traceability logs

Standard Operating Procedures

Methods pertaining to the conduct of this project

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# APPENDIX A

# **APPROVAL SIGNATURES**

**Project Title:** 

Biodegradation Screen Study for Telomer Alcohols

Client Project ID:

E01-0684

Contract Analytical Project Number: CA085

Report Revised by:

Mourad Rahi

Moradflel 11-7-02
Signature Warde 11/7/02

Laboratory Management:

Bruce E Warden

## Appendix B

# Final Quantitative Results for PFOA in samples, controls and blanks

**Table 1.** The measured PFOA concentrations in test cultures during the 16-day screening study for telomer alcohol aerobic biodegradability.

	Concentration (ng/mL) (Eluate 2)
Sample	PEOA
Day 0 biodegradation sample, CA085-0927-SA-031, E2	0.0
Day 1 biodegradation sample, CA085-0927-SA-033, E2	25 2
Day 2 biodegradation sample, CA085-0927-SA-035, E2	65.O
Day 5 biodegradation sample, CA085-0927-SA-037, E2	111.4
Day 7 biodegradation sample, CA085-0927-SA-039, E2	139.8
Day 14 biodegradation sample, CA085-0927-SA-041, E2	158.8
Day 16 biodegradation sample, CA085-0927-SA-043, E2	180,9

**Table 2.** The measured PFOA concentrations in the abiotic control cultures during the 16-day screening study for telomer alcohol aerobic biodegradability.

	Concentration (ng/mL) [Eluate 2]
Sample	PFOA
Day 0 no sludge control, CA085-0927-SA-053, E2	0.0
Day 1 no sludge control, CA085-0927-SA-055, E2	0,0
Day 2 no studge control, CA085-0927-SA-056, E2	0.0
Day 5 no studge control, CA085-0927-SA-059, E2	0.0
Cay 7 no studge comrol, CA085-0927-SA-060, E2	0.0
Day 14 no sludge control, CA085-0927-SA-062 E2	0.0
Day 16 no sludge control, CA085-0927-SA-064, E2	0.0

**Table 3.** The final data results for sludge blanks during the 16-day screening study for telomer alcohol aerobic biodegradability. PFOA was not detected in any of the sludge blanks.

	Concentration (ng/mt.) [Eluate 2]
Sample	PFOA
Day 8 studge blank, CA085-0927-SA-045, E2	0.0
Day 1 sludge blank, CA085-0927-SA-048, E2	0.0
Day 2 sludge blank, CA085-0927-SA-047, E2	0.0
Day 5 sludge blank, CA085-0927-SA-048, E2	0.0
Day 7 sludge blank, CA085-0927-SA-049, E2	0.0
Day 14 studge blank, CA085-0927-SA-050, E2	0.0
Day 18 sludge blank, CA085-0927-SA-051, E2	0.0

# Appendix C

# Integrated Chromatogram Peak Area Data for telomer alcohols

**Table 1.** Integrated LC/MS peak area raw data for the parent telomer alcohols substrates. Data acquired from sequence run CA085\_092701a.spl as SIR data.

Sample		Telomer Alcohol Integrated Peak Areas								
	File ID	C6	C8	C10	C12	C14	C15	SUM		
Oay 0 biodegradation sample, CA085-0927-SA-031, E2	CA085_092701a_008	3503	21704	28940	18446	5558	1326	1326		
Cay 1 biodegradation sample, CA085-0927-SA-033, E2	CA085_092701a_009	ND	91	1309	8633	4768	1340	1340		
Day 2 biodegradation sample, CA085-0927-SA-035, E2	CA085_092701a_010	ND	268	282	6677	4483	1213	1213		
Day 5 biodegradation sample. CA085-0927-SA-037, E2	CA085_092701a_011	12	690	103	3070	3701	1140	1140		
Day 7 biodegradation sample, CA085-0927-SA-039, E2	CA085_092701a_012	NΩ	631	ND	1263	3248	1343	1343		
Day 14 biodegradation sample, CA085-0927-SA-041, E2	CA085 092701a 013	ND	594	NO.	870	1324	1093	1093		
Day 18 biodegradation sample, CA085-0927-SA-043, E2	CA085_092701a_014	ΝD	643	117	685	1278	1137	1137		
Day 0 no sludge control. CA085-0927-SA-053, E2	CA085_092701a_016	5072	30841	22594	13734	5014	1492	1492		
Day I no sludge control, CA085-0927-SA-055, EZ	CA085 092701a_017	4891	31305	20373	11273	4712	1477	1477		
Day 2 no sludge control, CA085-0927-SA-056, E2	CA085_092701a_018	5198	31862	24414	13866	5379	1462	1482		
Day 5 no sludge control, CA085-0927-SA-059, E2	CA085_092701a_019	4383	27486	22470	12856	4941	1393	1393		
Day 7 no sludge control, CA085-0927-SA-060, E2	CA085_092701a_020	3515	21450	13821	10429	4252	1095	1095		
Day 14 no sludge control, CAO85-0927-SA-062 E2	CA085_092701a_021	3319	21284	14366	10226	4992	1265	1265		
Day 16 no studge control, CAO85-0927-SA-064, 62	CA085_092701a_022	4602	27329	17948	12215	5230	1476	1478		
Day 0 sludge blank, CA085-0927-SA-045, E2	CA085_092701a_044	ND	47	ND	ND	ND	ND	NO		
Day 1 sludge blank, CA085-0927-SA-046, E2	CA085_092701a_045	NO	ND	ND	ND .	NO	ND	NO		
Day 2 sludge blank, CA085-0927-SA-047, E2	CA085_092701a_046	ND	NO	ND	ND ·	NO	ND	ND		
Day 5 sludge blank, CA085-0927-SA-048, E2	CA085_092701a_047	ND	NO	ND	ND	ND	ND	ND		
Day 7 sludge blank, CA065-0927-SA-049, E2	CA085_092701a_048	37	162	ND	ND	ON	ND	NO		
Day 14 sludge blank, CA085-0927-SA-050, E2	CA085_092701a_049	ND	211	ND	מא	No	ND	ND		
Day 16 sludge blank, CA085-0927-SA-051, E2	CA085 092701a 050	17	68	ND	269	NO	ND	NO		

**Table 2.** Percentage of integrated peak area remaining at different time points with respect to the peak area at time zero for their respective culture types. Near complete degradation of several telomer alcohols (TA) was observed in test cultures only. Near 90% remained in control cultures for all except C<sub>10</sub> TA, which was 79.4%.

Sample Description		% TA Remaining versus Day 0						
		C <sub>6</sub> TA	C, TA	C <sub>10</sub> TA	C <sub>12</sub> TA	C <sub>14</sub> TA	C <sub>16</sub> TA	
Day 0 biodegradation sample, CA085-0927-SA-031, E2	0	100.0%	100.0%	100.0%	100.0%	100.0%	100.0%	
Day 1 biodegradation sample, CA085-0927-SA-033, E2	1	0.0%	0.4%	4.5%	46.8%	85.8%	101.1%	
Day 2 biodegradation sample, CA085-0927-SA-035, E2	2	0.0%	1.2%	0.9%	36.2%	80.7%	91.5%	
Day 5 biodegradation sample, CA085-0927-SA-037, E2	5	0.3%	3.2%	0.4%	16.6%	66.6%	86.0%	
Day 7 biodegradation sample, CA085-0927-SA-039, E2	7	0.0%	2.9%	0.0%	6.8%	58.4%	101.3%	
Day 14 biodegradation sample, CA085-0927-SA-041, E2	14	0.0%	2.7%	0.0%	3.6%	23.8%	82.4%	
Day 16 biodegradation sample, CA085-0927-SA-043, E2	16	0.0%	3.0%	0.4%	3.7%	23.0%	85.7%	
Day 0 no sludge control, CA085-0927-SA-053, E2	0	100.0%	100.0%	100.0%	100.0%	100.0%	100.0%	
Day 1 no sludge control, CA085-0927-SA-055, E2	1	96.4%	101.5%	90.2%	82.1%	94.0%	99.0%	
Day 2 no sludge control, CA085-0927-SA-056, E2	2	102.5%	103.3%	108.1%	101.0%	107.3%	98.0%	
Day 5 no sludge control, CA085-0927-SA-059, E2	5	86.4%	89.1%	99.5%	93.6%	98.5%	93.4%	
Day 7 no sludge control, CA085-0927-SA-060, E2	7	69.3%	69.6%	60.3%	75.9%	84.8%	73.4%	
Day 14 no sludge control, CA085-0927-SA-062 E2	14	65.4%	69.0%	63.6%	74.5%	99.6%	84.8%	
Day 16 no sludge control, CA085-0927-SA-064, E2	16	90.7%	88.6%	79.4%	88.9%	104.3%	99.1%	

# Appendix D

# Integrated Peak Area Data for Perfluorinated Acid End Products

**Table 1.** Integrated LC/MS peak area raw data for the end products formed. Data acquired from sequence run CA085\_092701a.spl as SIR data.

	- · · · · · · · · · · · · · · · · · · ·	face and Deep Area							
<u> </u>		integrated Peak Areas							
Sample.	File ID %	Perfluoro- pentanoate	Perfluoro- hexanoate	Perfluoro- heptanoate	Perfluoro- octanoate	Perfluoro- nonzanoate	Perfluoro- decanoate	Perfluoro- undecanoate	Perfluoro- dodecanoate
Day 0 bioxlegradation sample, CA085-0927-SA-031, E2	CA085_092701a_008	276	3576	12127	2715	8818	253	1559	375
Day I bindingradation sample, CA085-0927-SA-033, E2	CA065_092701a_009	13796	46183	15484	41821	5386	8428	1281	395
Day 2 biodigradation sample, CA085-0927-SA-035, E2	CA085_0927013_010	48384	127207	25841	88508	8267	17566	1378	1192
Day 5 biodingradation sample, CA085-0927-SA-037, E2	CA085_092701a_011	68648	204221	35747	140409	9375	44419	1641	2885 .
Day 7 biodegradation sample, CA085-0927-SA-039, E2	CA085_092701a_012	72870	210569	43227	170923	13751	75377	2844	5047
Day 14 biodegradation sample, CA085-0927-SA-041, E2	CA085_092701a_013	78978	221543	47751	190903	15382	92188	3071	11078
Day 18 biodegradation sample, CA085-0927-SA-043, E2	CA085_092701a_014	79826	215870	47315	192075	15889	96834	3278	11603
Day 0 no studge control, CA085-0927-SA-053, E2	CA085_092701a_018	193	861	9244	1057	4382	184	1283	49
Day 1 no studge control, CA085-0927-SA-055, E2	CA065_092701a_017	288	1722	16885	1695	5516	233	1692	51
Day 2 no sludge control, CA085-0927-SA-058, E2	CA085_092701a_018	351	1437	17394	1649	5958	164	1178	21
Day 5 no studge control, CA085-0927-SA-059, E2	CA085_092701a_019	475	1744	17948	1683	7420	278	1928	61
Day 7 no sludge control, CA085-0927-SA-060, E2	CA085_092701a_020	457	1643	15212	1583	8328	173	1015	25
Day 14 no sludge control, CA085-0927-SA-062 E2	CA085_092701a_021	581	1731	18978	1680	7129	214	1675	68
Day 16 no sludge control, CA085-0927-SA-064, E2	CA065_092701a_022	811	1851	17910	1837	7203	233	1342	34
Day 0 sludge blank, CA085-0927-SA-045, E2	CA085_092701a_044	618	525	31	117	18	0	0	3
Day I sludge blank, CA085-0927-SA-046, E2	CA085_092701a_045	683	152	76	166	262	0	0	- o
Day 2 sludge blank, CA085-0927-SA-047, E2	CA085_092701a_048	685	510	51	228	722	0	O	48
Day 5 sludge blank, CA085-0927-SA-048, E2	CA085_092701a_047	872	<b>J20</b>	86	223	392	0	0	Ð
Day 7 sludge blank, CA085-0927-SA-049, E2	CA085_092701a_048	737	248	82	581	247	0	24	42
Day 14 sludge blank, CA085-0927-SA-050, E2	CA085_092701a_049	876	237	86	731	314	D	24	29
Day 16 sludge blank, CA085-0927-SA-051, E2	CA085_092701a_050	634	285	81	645	397	0	21	0

# Appendix E

# Integrated Peak Area Data for Transiently formed Polyfluorinated Acid Products

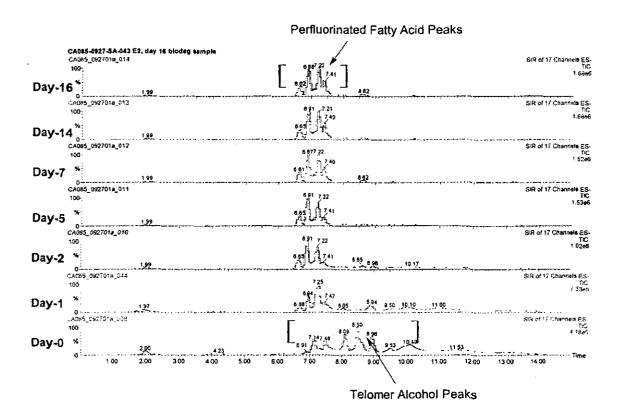
**Table 1.** Integrated LC/MS peak area data for transient products. Data from sequence CA085\_092401b.spl as full scan MS data from *m/z* 100 to *m/z* 1000.

		Integrated Peak Areas						
and the second		2H- perfluoro-2-	2H- perfluoro-2-	2H- perfluoro-2-	1H,1H- perfluoro-	1H,1H- perfluoro-	1H,1H- perfluoro-	
Sample	File ID	octenoate	decenoate	dodecenoate	octanoate	decanoate	dodecanoate	
Day 0 biodegradation sample, CA065-0927-SA-031, E2	CA085_092401b_008	1040	ND	ND	3085	211	ND.	
Day 1 biodegradation sample, CA085-0927-SA-033, E2	CA085_092401b_009	3429	760	249	15825	7065	958	
Day 2 biodegradation sample, CADS5-0927-SA-035, E2	CA085_0924015_010	223	ND	СN	8020	1802	263	
Day 5 biodegradation sample, CA085-0927-SA-037, E2	CA085_092401b_011	ND .	NO	NO	238	ND	ND	
Day 7 biodegradation sample, CA085-0927-SA-039, E2	CA085_092401b_012	ND	ND	ND	157	ND	ND	
Day 14 biodegradation sample, CA085-0927-SA-041, E2	CA085_092401b_013	ND	ND	ND	86	ND	ND	
Day 16 biodegradation sample, CA065-0927-SA-043, E2	CA085_092401b_014	ND	ND	NO	128	ND	ND	
Day 0 no sludge control, CA085-0927-SA-053, E2	CA065_092401b_616	ND	ND	ND OA	ND	ND	ND	
Oay 1 no sludge control, CA085-0927-SA-055, E2	CA085_092401b_017	ND	D	9	8	סא	ND	
Day 2 no siudge control, CA085-0927-SA-066, E2	CA085_092401b_018	ND	סא	ND	NO	NO	NO.	
Day 5 no studge control, CA085-0927-SA-059, E2	CA085_092401b_019	, ND	NO	ND	ND	ИĎ	ND	
Day 7 no sludge control, CA085-0927-SA-060, E2	CA085_092401b_020	ND	ND.	ND	NO	ND	ND	
Day 14 no sludge control, CA065-0927-SA-062 E2	CA085 092401b 021	ND	ND	ND	ND	ND	ND	
Day 18 no skudge comirot, CA065-0927-SA-064, E2	CA085_092401b_022	ND	ND	ND	ND	ND	ND	
Day 0 sludge blank, CA085-0927-SA-045, E2	CA085_092401b_030	ND	NO	NO	ND	ND	סא	
Oay 1 sludge blank, CA085-0927-SA-048, E2	CA085_092401b_031	ND	ND	ND	NO	ND	ND	
Day 2 sludge blank, CA085-0927-SA-047, E2	CA085_0924016_032	NĐ	ND	ND	ND	NO	ND	
Day 5 studge blank, CA085-0927-SA-048, E2	CA085_092401b_033	NĐ	ND	ND	ND ND	ND	ND	
Day 7 studge blank, CA085-0927-SA-049, E2	CA085 092401b 034	ND	ND	ND	ND	ND	ND	
Day 14 sludge blank, CA085-0927-SA-050, E2	CA085_092401b_035	ND	ND	ND	ND CM	ND	ND	
Day 16 sludge blank, CA085-0927-SA-051, E2	CA085_092401b_038	ND	ND	ND	ND	ND	ND	

# Appendix F

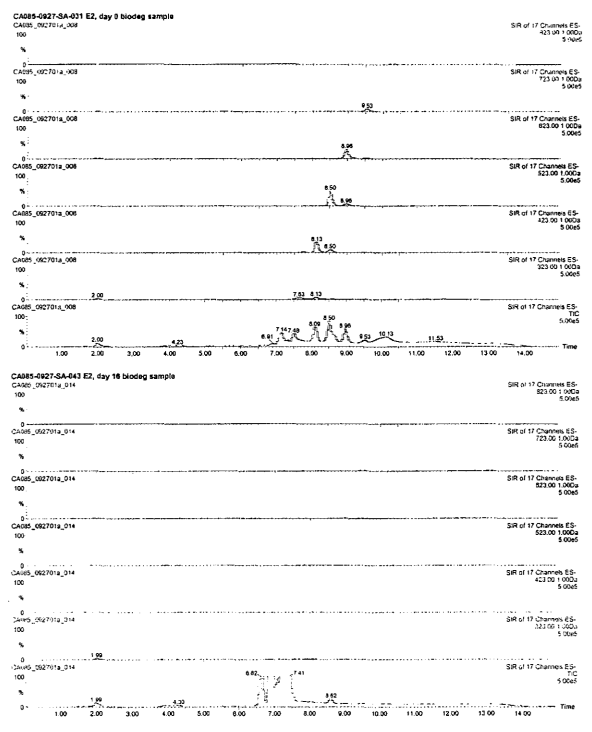
Typical Total Ion Chromatograms for SIR Data Showing Telomer Alcohols and Polyfluorinated Fatty Acid Products from Day-0 to Day-16 for HPLC/MS Analysis of Biodegradation Culture SPE eluate 2.

SIR of 17 anions: m/z 263, 313, 363, 413, 463, 513, 563, and 613 for perfluorinated acids; m/z 377, 477 and 577 for polyfluorinated acids; and m/z 323, 423, 523, 623, 723, and 823 for telomer alcohols.



## Appendix G

# SIR data Showing Ions for Telomer Alcohols at Day-0 and Day-16 from Analysis of the Biodegradation Culture SPE eluate 2.

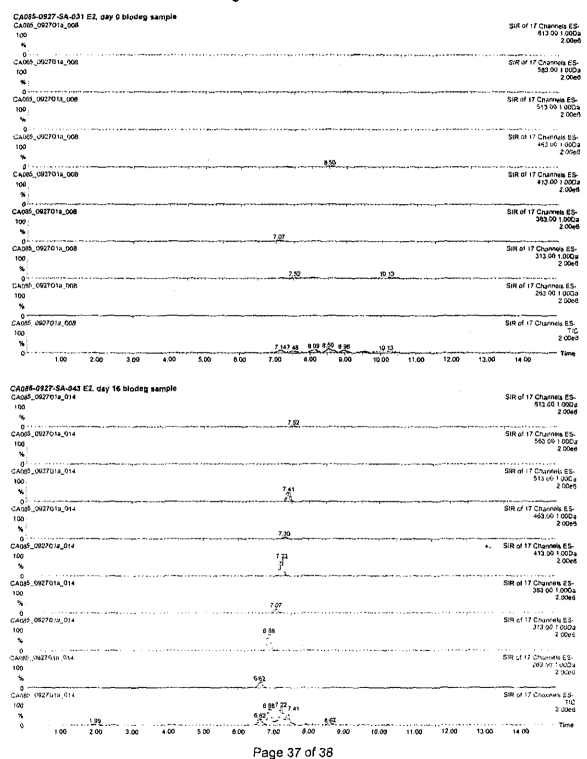


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# Appendix H

# SIR data Showing Ions for Perfluorinated Fatty Acids at Day-0 and Day-16 from Analysis of the Biodegradation Culture SPE eluate 2.

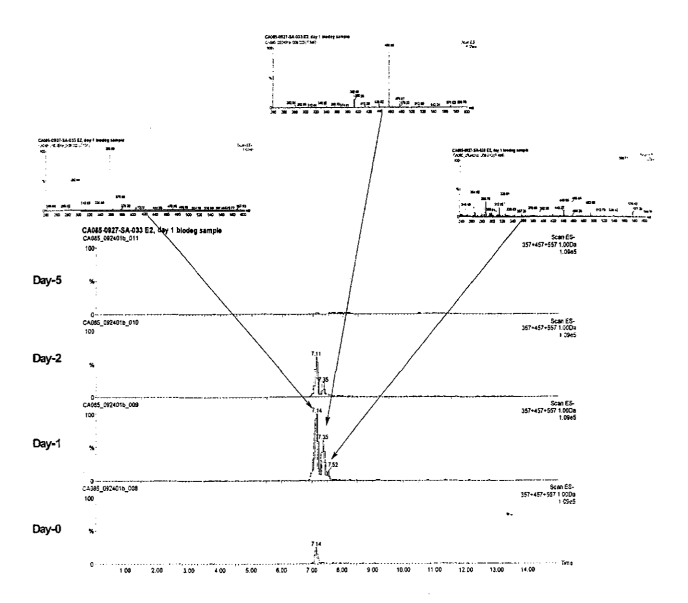


Pace Project CA085: "Biodegradation Screen Study for Telomer Alcohols" –Revision 1 Study Director: Cleston C. Lange, Ph.D. Pace Analytical Services, Bio-Analytical Services Group

Wednesday, November 06, 2002

# Appendix I

Chromatograms of Extracted ions m/z 377, 477, and 577 and Mass Spectra for Each Peak for Polyfluorinated Acid Intermediates Formed Transiently in Biodegradation Cultures at Day-0, Day-1, Day-2 and Day-5. Suspected β-Oxidation olefinic acid intermediates.



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